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Time-lapse scanning surface plasmon microscopy of living adherent cells with a radially polarized beam

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We report on a fibered high-resolution scanning surface plasmon microscope for long term imaging of living adherent cells. The coupling of a high numerical aperture objective lens and a fibered heterodyne interferometer enhances both the sensitivity and the long term stability of this microscope, allowing for time-lapse recording over several days. The diffraction limit is reached with a radially polarized illumination beam. Adherence and motility of living C2C12 myoblast cells are followed for 50 h, revealing that the dynamics of these cells change after 10 h. This plasmon enhanced evanescent wave microscopy is particularly suited for investigating cell adhesion, since it can not only be performed without staining of the sample but it can also capture in real time the exchange of extracellular matrix elements between the substrate and the cells.

OCIS codes: (240.6680) Surface plasmons; (180.5810) Scanning microscopy; (170.3880) Medical and biological imaging; (120.3180) Interferometry; (060.2310) Fiber optics; (110.4280) Noise in imaging systems.

1. INTRODUCTION

During the past decade, the evidence has become undeniable that in adhesion related processes the behavior of cells is not only controlled by biochemical cues but also involves many physical determinants like their structural and dynamical properties driven by molecular motors. The adhesion of living cells has an impact on their migration, division or death, differentiation, and surrounding tissue development [1]. It is governed by molecular interactions of the cytoskeleton (CSK) with the extracellular matrix (ECM) via signal transduction receptors (integrins and discoidin domain tyrosine kinase receptors) [2,3] and cytokine exchanges (growth factors) [4]. Integrin mediated adhesion is a complex cascade of biochemical and biomechanical events, for instance integrin binding to fibronectin (Fn) involves a conformational change (activation) of its Fn receptor that results in a mechanical coupling [5]. Bound Fns then rapidly associate with the actin CSK and cluster together, giving rise to adhesion complexes that contain cytoplasmic structural proteins such as vinculin and talin, and signaling molecules like focal adhesion kinase (FAK), Ras, and Src

[6,7]. Thanks to bidirectional mechanochemical mechanisms and their leading edge protrusions (lamellipodia and filopodia), living cells sense the stiffness of their environment and retroactively adjust their spreading, contraction, and migration [8]. Once driven to adhesion, they can choose among different types of adhesion interactions, mediated by proteins [9]. These include focal adhesions, close contacts, and ECM contacts (fibrillar adhesion). The strongest of these are focal adhesions, which have a tight contact within a distance of only 10–20 nm to the substrate. Close contacts often surround focal adhesions, but have a larger gap distance of about 30–50 nm. ECM contacts are long range adhesions with distances greater than 100 nm between the cell and the substrate. We choose in this work C2C12 myoblast cells for characterizing their adhesion and migration on gold surfaces. C2C12 is an immortalized mouse myoblast cell line that readily proliferates in high serum conditions, and differentiates and fuses in low serum conditions. It is a very useful system to study aspects of myogenesis, metabolism, and muscle biology [10]. C2C12 myoblast cell adhesion and motility are critical for their myogenesis.

During differentiation, these cells become more fusiform and aligned, exhibiting more intercellular contacts and cell membrane fusions leading to elongated contractile syncytia multinucleated myotubes [10].

High resolution characterization of living cell substrate interaction was achieved for the first time to our knowledge by total internal reflection microscopic (TIRM) observation of the cell substratum interface [11,12]. Combining TIRM and interferometric methods, interference reflection microscopes (IRMs) improved their sensitivity, allowing quantitative estimation of the cell substrate gap distances and, importantly, released the sample preparation from harmful staining procedures [13–15]. Due to its high sensitivity to refractive index (RI) changes at metal dielectric interfaces, surface plasmon resonance (SPR) was first introduced in the biology community as a biosensor for probing biochemical interactions and kinetics [16]. This optical technique was adapted subsequently for SPR imaging (SPRi) of cellular adhesion. Exploiting the amplification of surface plasmon polaritons (SPPs) [17] occurring on a metal dielectric interface, living cell SPRi was shown to provide a reliable estimate of cell surface distances at focal adhesion sites and close contacts [18]. SPRi was also used for long term imaging of protein salting out and cell substrate interactions [19,20]. Prism coupled SPRi was limited by the lateral propagation of SPPs, making cell details inaccessible. The introduction of high numerical aperture (NA) objective lenses [21–23] raised SPRi to the level of IRM and TIRM, up to what is called nowadays high resolution surface plasmon microscopy (HRSPM). Objective coupled SPPs are localized SPPs. Enhancement of the sensitivity and resolution of HRSPMs was improved in scanning versions of these microscopes by introducing interferometers [24,25] and confocal imaging [26–28]. Wide field (nonscanning) devices were also developed with (i) an annulus to confine the incoming light angles around the SPR angle [29] and (ii) a TIRM for focusing the light beam in the back focal plane of the objective lens at the SPR angle [30]. For both methods, the light reflected back from the sample was imaged on a 2D camera. HRSPM has been further applied to observe microstructures [24,31], micro and nanoparticles [23,32–34], bilipid layers [35,36], self assembled monolayers [30], and biomolecular binding events [27,37,38]. Cellular observation with HRSPM was performed on fixed cells in air [39–42] and in a liquid medium [43–45]. However, only a few experiments were carried out on living cells [46–49]. In Ref. [48], the strength of adhesion of single SH EP1 epithelial cells with collagen was measured as a function of the osmotic pressure using a TIRF SPRM configuration. However, full HRSPM was not exploited and cellular observations were limited in time to a few minutes. Lately, better resolution was obtained with a scanning HRSPM and a radially polarized light by Toma *et al.*, [49] and the cell substrate gaps were estimated for human embryonic kidney (HEK293) cells and poly L lysine coated gold surfaces, with bare gold as a control.

In this paper, as suggested by Somekh's group [24,25], we use a scanning IRM HRSPM to observe living mouse myoblast cells adherent on bare gold, with long term (50 h) time lapse video recording. We actually improve this technique by introducing a radially polarized illumination and a fibered

interferometer, both leading to better resolution and stability. We show that the resolution reaches the theoretical diffraction limit and that several days of recording can be achieved with this stabilized and compact setup. To the best of our knowledge, this is the first time that a scanning HRSPM (or a scanning surface plasmon microscope SSPM) is used for long term cell observation in a liquid medium. In the first part of the manuscript, we review all technical issues that contribute to a correct interpretation of the images produced by this optical device. The experimental setup is described with special attention to the fully fibered interferometer. The operating conditions and the contrast mechanism of this scanning HRSPM are detailed. The long term microscope stability is discussed and the resolution and penetration depth of the microscope are characterized. In the second part, we report on long term (48 h) imaging of living mouse myoblast cells. In particular, we reveal and quantify changes in the cell dynamics and their adhesion properties.

2. SSPM IN LIQUID

A. Experimental Setup

The SSPM principle is sketched in Fig. 1(a). SPPs are excited and confined with a 1.57 NA objective lens (Zeiss). When the objective lens is focused beyond the gold film ($z > 0$), into the liquid medium, SPPs propagate on the gold/liquid interface symmetrically toward the objective axis and, by leakage radiation (LR), a portion of them return to the back focal plane of the objective lens. The following two kinds of rays contribute to the back reflected light: the LR rays that have excited SPPs and the other rays that have not excited SPPs. Between these two kinds of rays, there is a phase difference due to defocus and

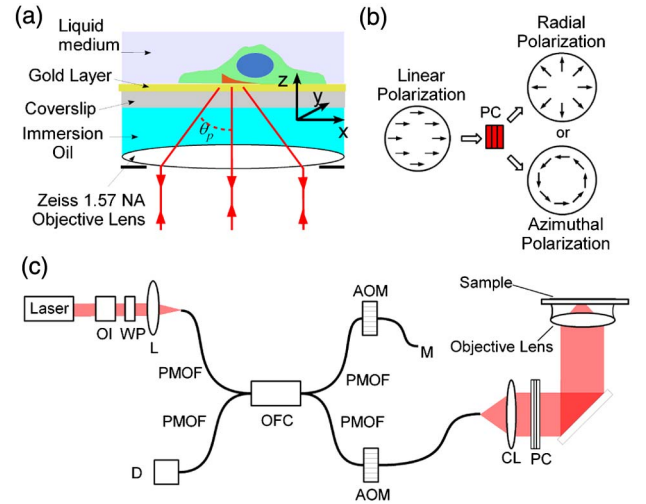


Fig. 1. Scanning surface plasmon microscope setup. (a) Sample configuration with the SPP excitation through a 1.57 NA (Carl Zeiss Microscopy GmbH) objective lens. (b) Spatial distribution of the polarization at the entrance of the objective lens aperture. The polarization converter (PC) switches between a radially and azimuthally polarized beam. (c) Microscope setup with the fully fibered interferometer. OI: optical isolator, WP: half wavelength waveplates, L: lens, PMOF: polarization maintaining optical fiber, OFC: optical fiber coupler, AOM: acousto optics modulator, D: detector, CL: collimating lens.

SPP propagation. This phase difference provides information on the SPP excitation conditions in the focused region and is accessed by an interferometric detection method. As SPPs are excited only with **p** polarization, the illumination of the objective lens aperture with a linearly polarized beam does not allow efficient excitation of SSPs. Actually, for a polar SPR angle θ_p [Fig. 1(a)], only a limited range of azimuthal angle φ contributes to the SPPs excitation. Given the radial geometry of the objective lens, all azimuthal angles can excite the SPPs [33,40,43] with a radially polarized beam. In this case, the polarization is purely **p**. For this purpose, we have introduced a liquid crystal cell polarization converter (PC) [50] to convert a linearly polarized beam into a radially polarized beam [Fig. 1(b)]. From this experimental setup, images with or without SPP excitation can be obtained by switching between radially and azimuthally polarized beams with the PC. The optical response of the microscope versus the defocus, known as the $V(z)$ curve, can be expressed in the integral form [24]

$$V(z) \propto \int_0^{2\pi} \int_0^{\theta_b} P^2(\theta) R(\theta) e^{i4\pi n_1 z \cos \theta / \lambda} \sin \theta d\theta d\varphi, \quad (1)$$

where $P(\theta)$ is the pupil function of the objective lens, θ and φ are the polar and azimuthal angles of the incoming light inside the objective lens, n_1 the refractive index of the coupling medium (the objective lens, the coverslip and the immersion oil), and $R(\theta)$ is the reflection coefficient of the sample. $R_p(\theta)$ [resp. $R_s(\theta)$] is obtained with a purely **p** (resp. **s**) polarization using a radially (resp. azimuthally) polarized beam.

As illustrated in Fig. 1(c), we have designed a fibered heterodyne interferometer allowing a better compaction of the experimental setup, together with some shortening of the time consuming optical alignment and adjustment steps. A Gaussian beam of a 1.5 mW stabilized red He Ne Laser is injected into the polarization maintaining optical fiber (PMOF). Inside the fiber, the light is separated in amplitude by a 50/50 optical fiber coupler (OFC). In the first arm, the light frequency is shifted by an acousto optical modulator (AOM) and is reflected at the end of the PMOF coated by a gold mirror. In the other arm, the light is frequency shifted by an AOM, exits the PMOF, is collimated by a 50 mm focal length lens (CL) and its polarization is converted by the PC before entering the high NA objective lens to reach the sample. Then the light is back reflected toward the CL and injected into the PMOF. The two path beams interfere and are detected by a phototube multiplier detector (D). Since, in SSPM, the polarization is a key parameter, in order to minimize depolarization inside the optical fibers we use PMOFs and weld all of the optical fiber components (PMOF, OFC, AOM) together. This leads to a polarization extinction ratio of 14 dB at the level of the PMOF output oriented toward the CL. The interferometric signal

$$I_{\text{int}}(x, y, z, t) = I_{\text{dc}} + I_{\text{ac}} = I_1 + I_2(x, y, z, t) + 2\sqrt{I_1 I_2}(x, y, z, t) \cos(2\pi f_m t + \psi(x, y, z, t)), \quad (2)$$

where I_1 (resp. I_2) is the optical power of the light coming from the first (resp. second) arm of the interferometer. I_{int} consists of a constant component I_{dc} and an alternative component I_{ac}

modulated at frequency f_m . A lock in amplifier (Signal Recovery model 7280) is used to extract from I_{ac} , the complex value:

$$I_{\text{SSPM}}(x, y, z, t) = 2A\sqrt{I_1 I_2}(x, y, z, t) e^{i\psi(x, y, z, t)}, \quad (3)$$

(in volts) at modulation frequency f_m , where $A = \text{GRgS}$ is the constant power to voltage conversion coefficient ($A \sim 5.3 \times 10^5$ V/W), defined as the product of the sensitivity $S = 0.05$ A/W of the phototube multiplier (D), the gain $g = 50$ of the detector, the resistance $R = 24$ k Ω of the current voltage converter, and the amplification gain $G = 12.5/\sqrt{2}$ of the lock in amplifier. A xyz piezoelectric stage (nPoint) is used to scan the sample in three spatial directions. This piezoelectric stage is mounted on an inverted microscope (IX71 Olympus). For a fixed xy position, a $V(z)$ curve is reconstructed from a z scanning:

$$V(z, t) = I_{\text{SSPM}}(z, t) = 2A\sqrt{I_1 I_2}(z, t) e^{i\psi(z, t)}. \quad (4)$$

Figure 2(a) shows in solid (resp. dashed) line the experimental $|V(z)|$ curves obtained for a 45 nm gold layer immersed in water for **p** (resp. **s**) polarization. The periodic oscillations visible for positive z defocus on the right of the central peak are a signature of SPP [24,25,51]. As expected for **s** polarization, the $|V(z)|$ curve decreases monotonously for positive z defocus; there are no oscillations. The complex reflectivity function $P^2(\theta)R(\theta)$ can be reconstructed by taking the inverse Fourier transform of the experimental $V(z)$ curve [Eq. (1)] [42,52,53] [Figs 2(b) and 2(c)]. The dashed curve in Fig. 2(b) can provide quantitative information on the beam profile at the objective lens aperture if we assume that in azimuthal polarization illumination $R_s(\theta)$ is nearly constant and equal to 1 and

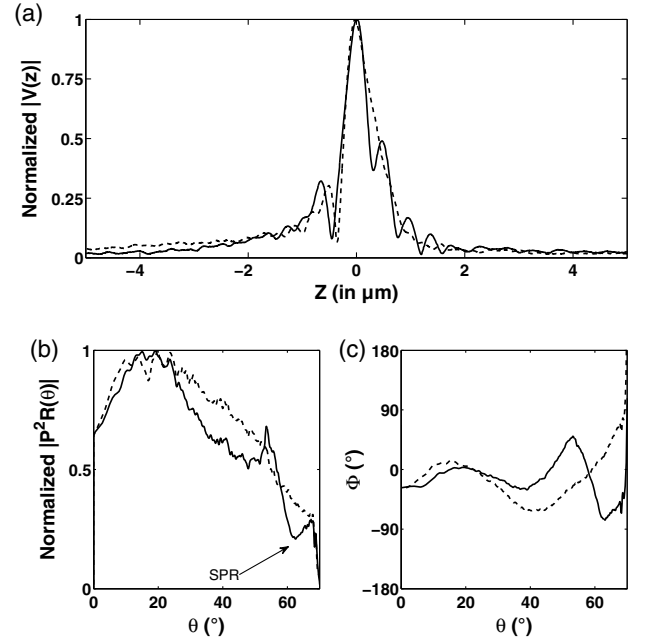


Fig. 2. (a) Experimental normalized $V(z)$ modulus curves for **p** polarization (solid line) and **s** polarization (dashed line) measured on a 45 nm gold layer in water. (b) and (c) represent the modulus and the phase of the reflectivity $P^2(\theta)R(\theta)$ versus the angle θ , computed from the experimental curves in (a), using Eq. (1) (see text).

that the beam profile has a radial symmetry for the angle φ . $\theta = 0^\circ$ and $\theta_{\max} = \arcsin(\text{NA}/n_1)$ correspond to the center and the border of the objective lens aperture, respectively. We can notice that the intensity of the beam decreases from the center toward the border of the objective lens aperture. This decrease corresponds to the Gaussian beam profile of the laser beam; the illumination beam diameter at the entrance of the objective lens is 9.6 mm, which is larger than the 6 mm diameter of the aperture. In the center of the beam, inside a disk with a diameter of 1.4 mm, the intensity decreases slightly due to a liquid crystal defect at the center of the polarization converter PC. As explained in a former work [32], the visibility of interference oscillations in the $V(z)$ curves depends on the ratio of intensities at the center and at the surface plasmon excitation angle. This ratio is nearly 2, which ensures a good weight for surface plasmon excitation and so a good visibility of the oscillations in the $V(z)$ curves. The SPR appears in the **p** polarization reflectivity curve at the angle $\theta_{\text{SPR}} = 62^\circ$. This SPR angle depends on the local refractive index of the medium in contact with the gold layer. Thus, as reported in Ref. [42], changes in the refractive index can be captured by measuring the local period of the $|V(z)|$ curve for positive z values. But, as for each (x, y) pixel the time required for $|V(z)|$ recording is ~ 10 s, it becomes quite a prohibitive task for a full (500×500) pixel matrix.

B. Image Reconstruction and Contrast Optimization

Imaging can be performed in a reasonable time (~ 50 s) as compared to the characteristic time of the dynamics of the object under study by fixing some z^* position selected from the $V(z)$ curves. For a given z^* position, a 500×500 pixel image can be recorded from 2D scanning along xy directions. For every pixel of the image, the raw intensity I_R is given by the modulus of I_{SSPM} :

$$I_R(x, y, t) = |I_{\text{SSPM}}(x, y, z^*, t)| = 2A\sqrt{I_1 I_2}(x, y, z^*, t). \quad (5)$$

Before each image acquisition, a single $V(z)$ curve is recorded at the first pixel of the image. This $V(z)$ curve is used as a reference curve to determine the exact position of the gold liquid medium interface (the value $z = 0$ corresponds to the position of the $V(z)$ maximum) and to normalize the raw image I_R by the maximum $V(z = 0)$. The normalized image I_N is then obtained as

$$I_N(x, y, t) = \frac{I_R(x, y, t)}{|V(z = 0)|}. \quad (6)$$

The contrast mechanism of SSPM imaging in a liquid medium is described by the $V(z)$ curves. When performing this imaging at a fixed scanning height z^* , the background intensity is not constant. These unwanted variations are due to some tilt between the xy scanning plane and the sample gold surface. In other words, the effective value of z^* is not maintained constant during xy scanning. We showed previously [54] that the z axis position can be adjusted in real time to keep the effective scanning height z^* constant. As compared to the method proposed in Ref. [54], here we will perform the tilt correction from the $|V(z)|$ curve obtained with an azimuthally polarized beam. As an illustration, we imaged in Fig. 3 a fixed C2C12 mouse myoblast cell maintained in a phosphate buffered saline solution

(PBS) (Appendix A). Note that changes in the $V(z)$ curve due to cells in an aqueous medium are expected to be smaller than in air since the refractive index drop between the cell and the culture medium is five to ten times smaller (0.03 to 0.06). We recorded 200 SSPM images for z^* , varying from $2 \mu\text{m}$ to $4 \mu\text{m}$, and for each radial and azimuthal polarization condition. A SSPM image I_N in radial polarization is shown in Fig. 3(a). Even if the tilt correction has been used during the xy scanning, it remains a small out of plane movement that we can estimate to be around 50 nm over the whole image ($110 \mu\text{m}$ size).

For each image $I_N(x, y, z^*)$, we have computed its probability density function (PDF) [black curve in Fig. 3(b)] and its cumulative distribution function (CDF) [Fig. 3(c)], from which a minimal I_N^- and maximal I_N^+ normalized intensity values have been determined at 5% and 95% of the CDF curve. $I_N^-(z^*)$ and $I_N^+(z^*)$ are plotted in Fig. 3(d). We notice that the two curves follow a typical $|V(z)|$ profile, which means that the residual tilt after correction becomes negligible as compared to the whole dynamics of the $|V(z)|$ curves. Note also that the tilt effect does not impact locally too much the intensity variation due to the presence of the object. Consequently, we will remove a background surface $I_B(x, y, t)$ [Fig. 3(e)] to each image to obtain the corrected image $I_C(x, y)$ [Fig. 3(f)]:

$$I_C(x, y, t) = I_N(x, y, t) - I_B(x, y, t). \quad (7)$$

For each z^* position, this background surface I_B is calculated by first fitting the image with a 2D polynomial function of order 2 in the x and y directions outside the cell and then by interpolating inside the cell. In the so corrected image I_C in

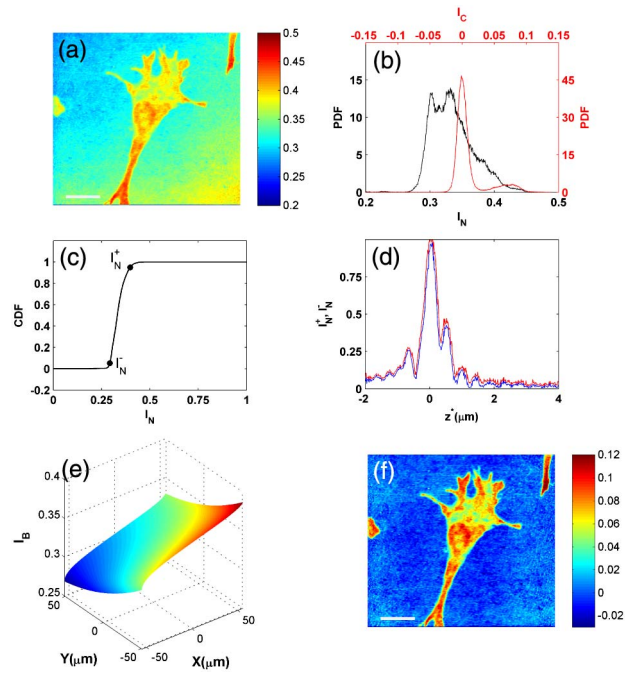


Fig. 3. (a) SSPM image I_N of a fixed C2C12 mouse myoblast cell in PBS. (b) Probability density function (PDF) of I_N (black) and I_C (red). (c) Cumulative distribution function (CDF) of I_N . (d) I_N (blue curve) and I_N^+ (red curve) versus z^* . (e) Background surface I_B (see text). (f) Corrected SSPM image $I_C = I_N - I_B$ [Eq. (7)]. The scale bar in (a) and (d) is $20 \mu\text{m}$.

Fig. 3(f), we can observe that the cell is easily distinguishable from the background. Interestingly, the PDF of the corrected image I_C [red curve in Fig. 3(b)] reveals the existence of a bimodal distribution as the signature of the presence of two populations that can be easily separated. The first one is centered at zero and corresponds to the image background (gold roughness, noise, and out of plane scanning), whereas the second distribution, centered around 0.075, is related to the cell adsorbed on the gold layer.

In Fig. 4, corrected SSPM images I_C are shown for radially and azimuthally polarized beams at positions $z^* = 0.39 \mu\text{m}$ and $z^* = 1.05 \mu\text{m}$, respectively. The cell is far less visible with an azimuthally polarized beam [Fig. 4(b)] than with a radially polarized beam [Fig. 4(a)]. To provide a better insight into the image formation with SSPM, let us investigate the evolution of the SSPM images versus the scanning height position z^* and the polarization illumination conditions. In Fig. 4(c), the $|V(z)|$ curves have been reconstructed from 200 SSPM images I_N for z^* between $-2 \mu\text{m}$ to $4 \mu\text{m}$, at pixels A and B shown in Fig. 4(a), for radial and azimuthal polarization illuminations. The $|V(z)|$ curves for the radially polarized beam present differences between the two positions whereas for the azimuthally polarized beam the two curves are almost superimposed.

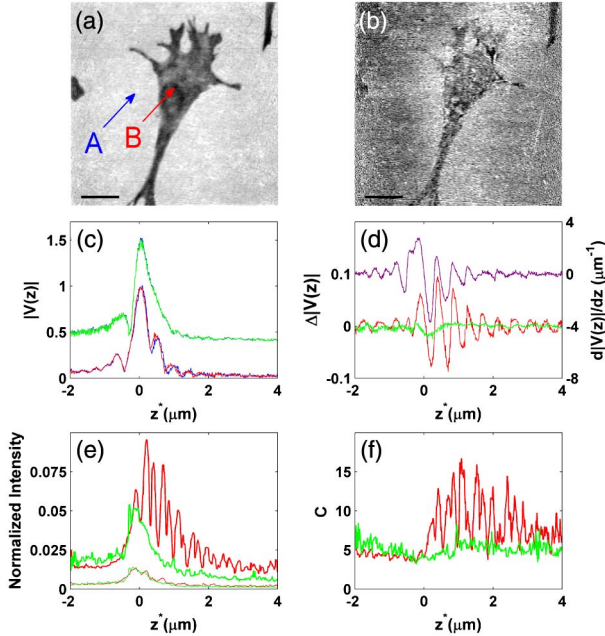


Fig. 4. (a) Corrected SSPM image I_C of the same fixed myoblast as in Fig. 3(a) obtained for radial polarization illumination at scanning position height $z^* = 0.39 \mu\text{m}$; the intensity scale is $[0.02; 0.12]$. (b) Same as in (a) for azimuthal polarization illumination at $z^* = 1.05 \mu\text{m}$, the intensity scale is $[0.006; 0.008]$. In (a) and (b), scale bars are $20 \mu\text{m}$. (c) Reconstructed $|V(z)|$ curves at pixels A (outside the cell) and B (inside the cell) from 200 images I_N , in radial polarization: A (blue) and B (red), and in azimuthal polarization: A (black) and B (green). For clarity, the $|V(z)|$ curves in azimuthal polarization are vertically shifted by 0.5. (d) $\Delta|V(z^*)|$ and $\frac{d|V(z^*)|}{dz}$ (purple). (e) Image dynamics $\Delta I_C(z^*)$ (thick lines) and rms roughness $\sigma_b(z^*)$ (thin lines). (f) Image contrast $C(z^*)$. In (d), (e), and (f), the red (resp. green) lines correspond to radially (resp. azimuthally) polarized beam.

The differences $\Delta|V(z)|$ between the two $V(z)$ curves, for radially and azimuthally polarized beams, are shown in Fig. 4(d). For radially polarized beams, $\Delta|V(z)|$ varies with z^* and oscillates between positive and negative values for positive defocus as the signature of some shift in the local period of $|V(z)|$ caused by refractive index variations. For azimuthally polarized beams, $\Delta|V(z)|$ is by far smaller than for radial polarization and does not show any oscillations. This illustrates the inefficiency of azimuthal polarization to characterize refractive index changes.

Let us now extend our single pixel analysis to the entire image I_C , and introduce the notions of image dynamics and contrast. For each image, we define its dynamics by $\Delta I_C = I_C^+ - I_C^-$, where I_C^+ and I_C^- correspond to the 95% and 5% CDF levels, respectively. As shown in Fig. 3(b), the PDF of I_C does not follow a normal distribution and the part of the histogram related to the cell has a smaller weight and is well separated from the background at higher intensity values. This argues in favor of using the CDF instead of the PDF standard deviation for computing I_C^+ and I_C^- . The image dynamics ΔI_C [Fig. 4(e)] evolves versus the scanning height position z^* with clear oscillations for radial polarization illumination (thick red line), whereas the curve obtained for azimuthal polarization illumination (thick green line) shows a single peak around $z^* = 0$ without oscillations and is always below the curve for radial polarization illumination. This underlines the amplification of the image dynamics for radially polarized beams as compared to azimuthally polarized beams. The amplification is supported by SPPs. To quantify the image contrast versus z^* and versus the polarization mode, the image dynamics ΔI_C must be compared to the background noise level for each image. The limit of detection of the SSPM in liquid is not ruled by the electronic noise (which is 0.0018 and can be lowered to 0.0009 if the frequency modulation f_m is increased from 140 to 1200 kHz), but by the background signal, mainly due to the gold roughness. The standard deviation σ_b of a bare gold area of the image I_C is computed for every z^* and is represented in Fig. 4(e) by a red (resp. green) thin line for radial (resp. azimuthal) polarization illumination. The background noise for both polarization conditions is z^* dependent since it is maximal near $z^* = 0$ and decreases with the defocalization z^* . We define the image contrast C as the ratio between the image dynamics ΔI_C and the background noise standard deviation σ_b for each scanning height position z^* :

$$C(z^*) = \Delta I_C(z^*) / \sigma_b(z^*). \quad (8)$$

$C(z^*)$ is shown in Fig. 4(f) for radially polarized (thick red line) and azimuthally polarized (thick green line) beams. The former $C(z^*)$ displays oscillations and values higher than 11 for a wide range of z^* values (between 0.39 and $2.9 \mu\text{m}$). The latter $C(z^*)$ is lower and much flatter and almost independent of z^* . These results show that SPPs are efficient in contrast amplification for z^* between 0.39 and $2.9 \mu\text{m}$. Between $z^* = 0$ and $0.39 \mu\text{m}$ and above $2.9 \mu\text{m}$, the image contrast produced by SPPs is hardly distinguishable from that obtained without SPPs. Interestingly, the z^* positions where $C(z^*)$ is locally maximum correspond to the extrema of $\Delta|V(z)|$ [red curves in Figs. 4(d) and 4(f)], which confirms that the contrast is strongly dependent on the shape of $|V(z)|$. However, the highest contrast

value is obtained for $z^* = 1.12 \mu\text{m}$ rather than for $z^* = 0.22 \mu\text{m}$, as expected from $\Delta|V(z^*)|$ and the image dynamics $\Delta I_C(z^*)$ curve in Figs. 4(d) and 4(e).

Experimentally, we need to select the value of z^* to perform the image xy scanning. In Fig. 4(f), we observe that the contrast $C(z^*)$ is strongly oscillating in the 0.39 and $2.9 \mu\text{m}$ z^* range, so that a 100 nm drift in z^* may dramatically decrease the contrast. We have also noticed that the positions for which the image contrast is locally maximum fluctuate from sample to sample over a range of 200 nm . These variations may be attributed to fluctuations in gold and coverslip thicknesses. Actually, thickness changes of the gold layer mainly entail modifications of the $|V(z)|$ oscillation amplitude, whereas mismatches between the real and optimal coverslip thicknesses introduce phase aberrations that shift the $|V(z)|$ oscillation positions. Since drawing similar $\Delta V(z^*)$ or $C(z^*)$ curves for each sample is prohibitively time consuming, a simpler and faster way to guess z^* positions with maximal contrast is to use the derivative $\frac{d|V(z)|}{dz}$ computed from one single $|V(z)|$ curve. As shown in Fig. 4(d), the $\Delta|V(z)|$ oscillations are strongly correlated to those observed in $\frac{d|V(z)|}{dz}$; the extrema fall at the same z^* positions within an error of $\sim 30 \text{ nm}$. Note that this comparison is possible because the local RI changes induce very little variations in the $|V(z)|$ curves.

C. Sensitivity and Limit of Detection

To complete our discussion of the choice of z^* for optimal contrast, we must also estimate the sensitivity S_{SSPM} and the detection limit in RI units (RIU) of the microscope at this position z^* when the refractive index n of the medium changes. The sensitivity S_{SSPM} can be computed from the value of $|V(z_m)|$ taken at the position z_m corresponding to a maxima of $\frac{d|V(z_m)|}{dz}$:

$$S_{\text{SSPM}} = \frac{\delta|V(z_m)|}{\delta n}. \quad (9)$$

Using a 4 layer model (Appendix C), the theoretical value $S_{\text{SSPM}}^{\text{theo}}$ was estimated at ~ 3.3 for RI drops in the interval $[10^{-4}, 6 \cdot 10^{-2}]$. The experimental $S_{\text{SSPM}}^{\text{exp}}$ was measured from mixtures of water and glycerol, with RIs calibrated with a refractometer, giving RI drops from 8.7×10^{-3} to 5.3×10^{-2} . The single point limit of detection of the SSPM microscope is actually given by the footprint $|V_{\text{sn}}(z)|$ of the shot noise on the $V(Z)$ curves. In heterodyne interferometry, the signal to shot noise ratio $|V(z)|/|V_{\text{sn}}(z)|$ is written as the ratio of the variance of the detector current $\langle i_{\text{ac}}^2 \rangle_t$ over the variance of the shot noise current $\langle i_{\text{sn}}^2 \rangle_t$

$$\frac{|V(z)|}{|V_{\text{sn}}(z)|} = \sqrt{\frac{\langle i_{\text{ac}}^2 \rangle_t}{\langle i_{\text{sn}}^2 \rangle_t}} = \sqrt{\frac{2\tau S}{e} \frac{I_1 I_2}{I_1 + I_2}} \simeq \sqrt{\frac{2\tau S}{e}} I_2, \quad (10)$$

where e is the electron charge and τ is the integration time of the lock in amplifier ($\tau = 100 \mu\text{s}$). For our experimental setup at $z_m = 1.12 \mu\text{m}$, the optical power $I_2 = I_1/270 = 50 \text{ nW}$ leads to $|V(z)|/|V_{\text{sn}}(z)| \sim 1.8 \times 10^3$. Hence, a shot noise level $|V_{\text{sn}}(z)| = 8.33 \times 10^{-5}$ on $V(z)$ curves gives a limit of detection of $\sim 5.6 \times 10^{-5}$ RIU. This theoretical limit could be further improved by a factor of 30 by (i) replacing the 50 mm focal

length lens by a shorter focal length lens system and adding a beam expander system for a better light coupling and (ii) replacing the photomultiplier tube detector by a photodiode (the quantum efficiency and so S could be enhanced by seven). Such modifications would lead to an ultimate theoretical limit of detection of 1.8×10^{-6} RIU for this SSPM technique.

In imaging mode, additional sources of noise are involved that lower the RI limit of detection. The gold layer roughness generates a background noise signal σ_b estimated at ~ 0.0035 , from the thin red curve on Fig. 4(e) at $z_m = 1.12 \mu\text{m}$. This value is identical to that obtained from a bare gold surface in water, meaning that cellular fragments that could deposit on the gold surface do not change drastically the SSPM signal fluctuations. From $S_{\text{SSPM}}^{\text{exp}}$, we compute the RI limit of detection: $\Delta n_{\text{lim}} = \sigma_b/S_{\text{SSPM}}^{\text{exp}} \sim 2.3 \times 10^{-3}$ RIU. Loison and Fort [55] estimated a detection limit of 4×10^{-4} RIU, which can be concluded as a six fold improvement. However, the RI drop detection limit is not independent of the thickness and the area of the considered dielectric layer, since the HRSPM microscope is sensing a whole volume in the evanescent field of the gold/dielectric interface. For comparing different experimental works, we thus need to compute $V_I = hA\Delta n$, where h is the thickness of the dielectric layer and A is the area of the sensed region (proportional to the PSF of the microscope). For a PSF of 181 nm (see Section 2.D), with the same 4 layer model as above (Appendix C), we get $\sigma_b = 0.0035$, leading to a volume detection limit of $\sim 4720 \text{ nm}^3$. In comparison, the volume detection limit obtained in Ref. [55] $\sim 15000 \text{ nm}^3$ is much less impressive. In the same line, Watanabe *et al.*, [35] used micro patterned phospholipid bilayers with a $4 \mu\text{m}$ diameter to conclude that their localized surface plasmon has a thickness resolution of 0.33 nm for an index drop of 0.113 , which would give a volume detection limit of 1080 nm^3 , which is four times smaller than what the SSPM device described here can afford. However, it must be noted that the computations in Ref. [35] were limited to small (micron size) structures with well defined bilipidic layers and did not take into account the background noise on large surface areas (as we do here). Finally, the HRSPM techniques in Refs [35,55] are only sensitive to SPR angle shift, whereas SSPM is also sensitive to SPR angle spreading coming from radiative leakage. Since gold layer roughness (i.e., local submicronic gold thickness variations) induces radiative leakage, the SSPM background noise is amplified and the RI detection limit is increased. Given that the SSPM background fluctuations can be scaled down by a factor of 3 to 4 by a surface treatment of the glass coverslip prior to gold sputtering [56], SSPM could nevertheless reach the same volume detection limit as other HRSPMs.

D. Stability for Time-lapse Imaging

The use of an interferometer automatically raises the issue of stability and reproducibility of the optical signal and consequently of the stability and reproducibility in SSPM imaging. There are four bottlenecks that should be discussed: (i) incorrect light retro injection through the CL inside the PMOF, (ii) errors in the distance z^* given by mechanical and thermal drifts of the sample, the objective lens and the microscope stand, (iii) out of plane movements produced by the xyz

piezoelectric stage in the z direction and positioning errors in xy directions, (iv) spurious reflected light that perturbs the interference signal I_{ac} . In this section, we discuss the stability of the SSPM with tentative estimation of the respective weights of these four kinds of instability.

Concerning thermal drift [bottleneck (ii)], an autofocus is performed at the beginning of each image, since the recorded $|V(z)|$ curve provides the exact position of the gold surface. The autofocus compensates drastically the thermal and mechanical drifts of the setup in the z direction. Actually, we have estimated a root mean square drift equal to 9.4 nm in z positioning for 27,183 recorded images (491 h recording). If we convert this spatial drift in terms of intensity I_N by multiplying the drift by the derivative of $|V(z)|$, we get an intensity drift ranging between 0.012 and 0.0188. The influence of instabilities of types (i) and (iv) can be studied from the reproducibility of $|V(z)|$ curves at one xy position on the surface. The instability (iv), due to spurious reflected light, can be computed from the $|V(z)|$ curves since it appears as a very slowly varying function of z on the real and imaginary parts of $I_{SSPM}(x, y, z, t)$ [Eq. (3)]. The fluctuation standard deviation due to spurious reflected light, when estimated over 175 min (100 $|V(z)|$ curves), is 0.025. After elimination of these fluctuations, the remaining $|V(z)|$ fluctuations are smaller than 0.015. From the 5 nm accuracy in the z positioning, we estimate the contribution to $|V(z)|$ of the type (iii) instability to be 0.0075 ± 0.0025 . The fluctuations due to the type (i) instability, i.e., light retro injection, produce an error of 0.0075 ± 0.0025 in $|V(z)|$. If these fluctuations are compared to the I_N value at $z^* \sim 0.4 \mu\text{m}$, they represent for instabilities i), ii), and iv) no more than 1.8, 1.8, and 6.3%, respectively. Consequently, the total fluctuations $\sim 9.9\%$ comes mainly from a spurious reflected light instability.

We have tested the overall imaging stability and reproducibility on 180 images of the fixed myoblast cell of Fig. 4, recorded for the same scanning height position $z^* = 0.39 \mu\text{m}$ (195 min). When considering the normalized images $I_N(x, y)$, we find that for each pixel of the image the temporal standard deviation of $\sigma_{I_N}(x, y)$ ranges between 0.019 and 0.0309, i.e., between 5 and 10% of the whole image time averaged normalized intensity $\langle I_N \rangle_t$. If we consider now the corrected image I_C , we get from the 180 images that inside the cell, the time standard deviation of $\sigma_{I_C}(x, y)$ is ~ 0.006 , as shown in Figs. 5(c) and 5(d) (the borders of the cells were not taken into account because of a drift in xy direction between images). This means that the errors induced by the instability of the interferometers are, in the worst case, at the same level as the background noise σ_b of the image (which is equal to 0.0069 at $z^* = 0.39 \mu\text{m}$). As shown in Figs. 5(e) and 5(f), the signal to noise ratio $\text{SNR}(x, y) = \langle I_C(x, y) \rangle_t / \sigma_{I_C}(x, y)$ (where $\langle I_C(x, y) \rangle_t$ is the time average of images I_C) inside the cell is as large as 10 to 30. Then, by inverting the SNR, we get that σ_{I_C} inside the cell is between 3.5 and 10% of the time averaged image intensity $\langle I_C \rangle_t$. This fluctuation estimate seems to be lower than the one extracted from the $V(z)$ curves. In fact, a significant amount of the fluctuations has been removed when performing the background subtraction to get the corrected intensity I_C [Eq. (7)]. This measurement of stability on fixed

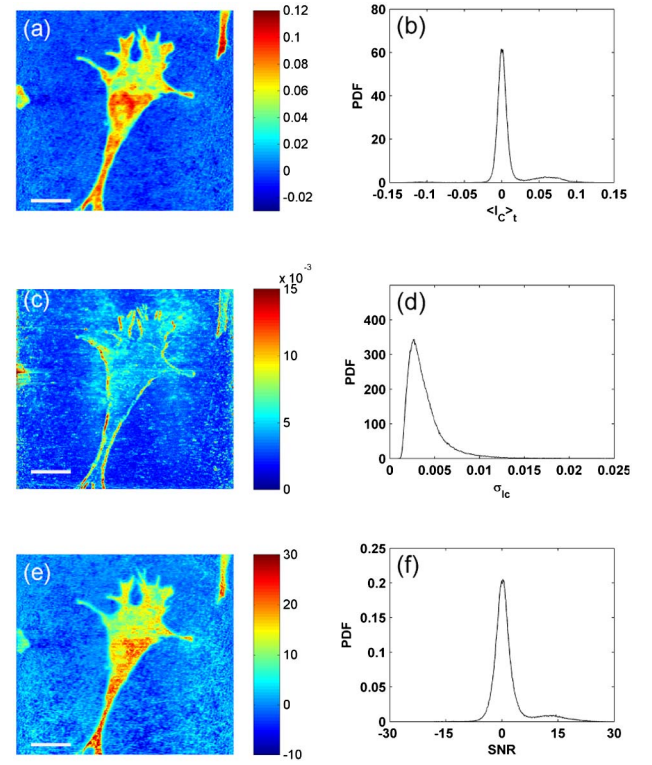


Fig. 5. (a) Time averaged $\langle I_C(x, y) \rangle_t$, over 180 images of a fixed myoblast cell, recorded over 195 min at $z^* = 0.39 \mu\text{m}$. (c) Corresponding standard deviation $\sigma_{I_C}(x, y)$. (e) Signal to noise ratio $\text{SNR}(x, y) = \langle I_C(x, y) \rangle_t / \sigma_{I_C}(x, y)$. (b), (d), and (f) correspond to the PDF of pixel values of the images shown in (a), (c), and (e), respectively.

cells confirms that the SSPM images are stable over a long period of time. This is the demonstration that the microscope with the heterodyne optical fiber interferometer provides a good stability. Another main advantage due the optical fiber design is the drastic minimization of optical alignment operations. For this microscope set up, no optical alignment or adjustment was needed for a period of 22 months, even if the microscope was used on a daily basis. Except for the coverslip, nothing else was changed on the microscope setup during this period. In summary, the stability of the heterodyne optical fiber interferometer opens the way for time lapse imaging of the dynamics of living cells.

E. Resolution and Penetration Depth

As shown in Fig. 4(a), the contour of the fixed myoblast cell is well resolved with submicronic resolution. As measured with a 50 nm gold particle, the experimental point spread function (PSF) of the SSPM in aqueous medium in radial polarization has a full width at half maximum (FWHM) of $\sim 180 \text{ nm}$ in x and y directions (Fig. 6). We conclude that the SSPM spatial resolution is limited by diffraction and is quite consistent with the theoretical value of 150 nm, obtained by computing the electromagnetic field near the focus of a radially polarized beam illuminating a 45 nm gold layer immersed in water (data not shown). The small difference between the experimental and

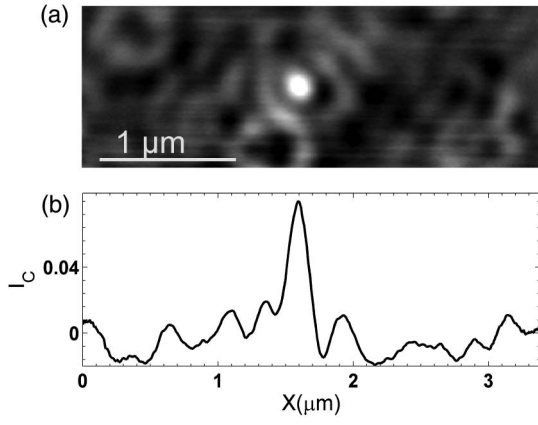


Fig. 6. (a) SSPM image I_C with a radial polarization illumination at $z^* = 0.425$ μm of a 50 nm gold particle in water. The intensity scale is [0.02; 0.06]. (b) Intensity profile I_C along a horizontal section crossing the particle in its center. The FWHM of the central peak is 181 nm.

theoretical values is certainly due to the aberrations introduced by the thickness mismatch of the coverslip.

SPPs probe the cell region that is inside the evanescent field. In the microscope, the image reconstruction is performed at a defocused position ($z^* > 0$), making the relation between the region probed by the SSPM and the SPP penetration depth not immediate, since the cell modifies locally the evanescent field penetration length. To estimate the depth of the probed region with the SSPM relative to the SPP penetration depth, we have imaged a 10 μm polymer sphere (polymethylsilsesquioxan Kobo diasphere KS 1000 sphere), with the SSPM at different positions z^* . Imaging has been performed in an ethanol/water (50/50%) medium in order to lower the RI drop between the object ($n = 1.41$) and the immersion medium ($n = 1.362$) to the typical RI drop of biological cells in their culture medium. The area probed with the SSPM [Figs. 7(b) and 7(d)] is by far smaller than the sphere size [Fig. 7(a)]. When moving away from the center of the particle, the sphere is no longer in contact with the surface and its separation distance increases [Fig. 7(c)]. The lateral position X_p where the particle is no longer detected with the SSPM provides an estimate of the depth of penetration $d_p = R \sqrt{R^2 - X_p^2}$, where R is the sphere radius [Fig. 7(c)]. To determine X_p from the SSPM intensity profile of the particle, two criteria can be used: X_p is estimated when the normalized intensity of the image (i) is divided by a factor ϵ or (ii) equal to the background noise standard deviation σ_b . As shown in Fig. 7(e), d_p evaluated using either criteria definitely increases as the defocus position z^* increases (from 100 to 300 nm for the former and from 180 to 600 nm for the latter). This is another difference from other TIRF based HRSPM [45] or scanning HRSPM [35]. In SSPM, the electromagnetic field at the interface is not a simple evanescent wave; part of the laser beam that does not undergo total reflection generates a focused beam above the gold surface whose position moves away from the gold interface when increasing the height position z^* . This explains the increase of d_p with z^* reported in Fig. 7(c). In the following application to

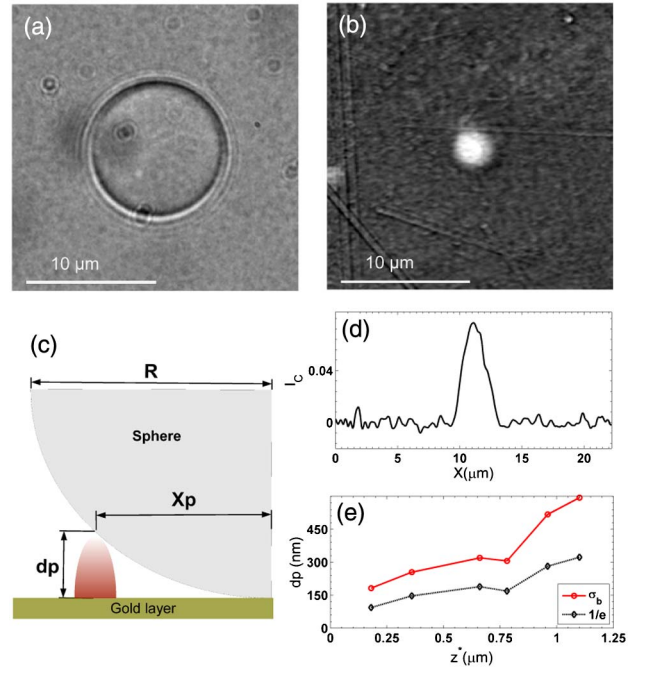


Fig. 7. Penetration depth of the SSPM probed with a 10 μm polymer bead immersed in a 50%/50% ethanol water medium probed by SSPM. (a) Transmission microscopy image. (b) SSPM image I_C with a radially polarized beam at scanning height position $z^* = 0.36$ μm of the same bead. (c) Sketch of the field probed by the SSPM. (d) Intensity profile I_C along a horizontal section crossing the particle in its center. (e) Evolution of the penetration depth d_p versus the scanning height position z^* using criteria (i) (black) and (ii) (red) (see text).

living cell imaging, we have chosen z^* at the first maximum of the derivative of $V(z)$ [Fig. 4(b)], between 350 and 500 nm corresponding to a penetration depth between 250 and 290 nm. We have made this choice in order to get the best contrast and to restrain the probed domain of biological cells only to the near field of SPPs.

3. LIVING CELL IMAGING

As a first application of SSPM to living cell imaging in liquid, we use again the C2C12 myoblast cell model under the cell culture and sample preparation conditions described in Appendix A. To get the best contrast, and to restrain the cell probed domain to the near field of SPPs, we will fix z^* at 0.4 μm, the first maximum of the derivative of $V(z)$ [Fig. 4(b)]. The living cells were observed with the SSPM at room temperature (24°C) during 50 h, with a time interval of 65 s.

A. SSPM Cell Imaging

From the time lapse video [Visualization 1](#) the inter and intra cellular dynamics are noteworthy. During the 50 h recording of the same zone of the coverslip, cellular migration, intercellular contacts, lamellipodial and filopodial dynamics, adhesion, detachment and attachment on the gold substrate are observable. In Figs. 8(a) 8(c) and 8(d) 8(f), two sequences of cell migration are reported, showing that different cells migrate over the same recorded area during 50 h. SSPM images show different

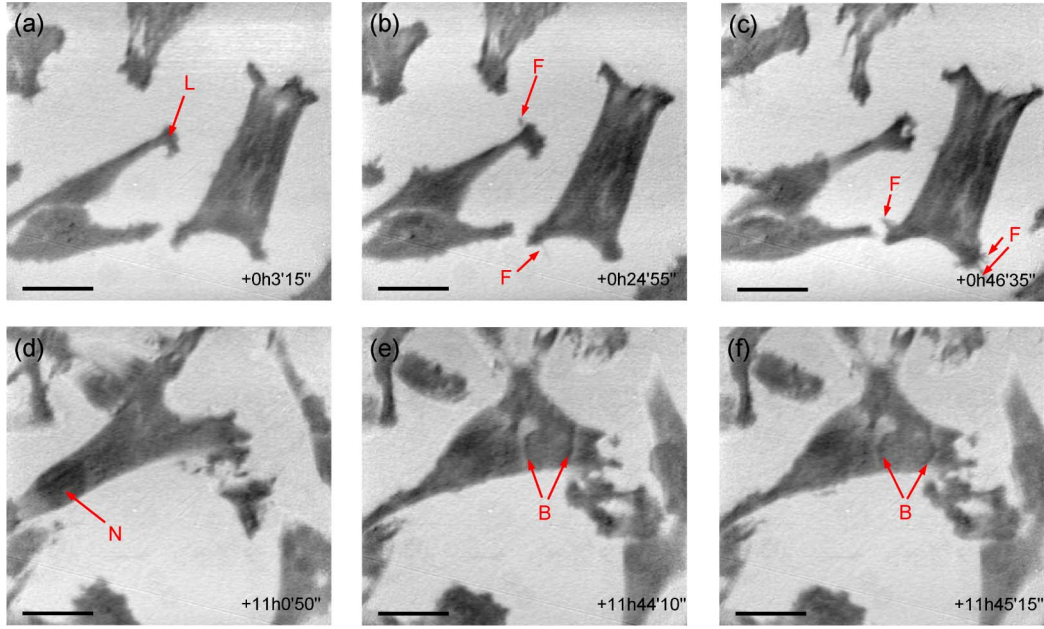


Fig. 8. SSPM images and time lapse video ([Visualization 1](#)) of living C2C12 cells during 50 h at scanning height position $z^* = 0.40 \mu\text{m}$. The scalar bar in all images is $20 \mu\text{m}$ and the intensity scale is $[0.02; 0.1]$. The arrows have the following meaning: filopodia (F), lamellipodia (L), nucleus (N) and fronts (B).

structures as filopodia [F arrows in (b) and (c)] and lamellipodia [L arrows in (a)]. The filopodia, which are few micrometers wide or less, are very dynamic; they explore the surface within a few micrometers distance and the same filopodia is not observable in more than two successive images ($\Delta t \sim 65 \text{ s}$), giving a protrusion retraction velocity larger than $3 \mu\text{m/s}$. Lamellipodia at cell extremities explore the gold surface, but also the neighboring cells; they show expansion and retraction during variable periods of time from minutes to more than an hour. During the time lapse video [Visualization 1](#), we can see that the whole cell bodies migrate, for example, the central cell in [Fig. 8\(d\)](#) moves from left to right in [Fig. 8\(f\)](#). Inside cells, very intense fronts with crescent forms propagate across the cells [B arrows in [Figs. 8\(e\)](#) and [8\(f\)](#)] during rather short periods of time (a few tens of minutes). These propagating fronts of locally higher RI volume correspond to either some parts of the cells that are denser near the surface or to some parts of the cells that are closer to the surface. In both interpretations, these propagating waves are likely to correspond to some actin network reorganization. Although the probed region inside the cells does not exceed 290 nm , the nucleus sometimes approaches close to the surface [N arrow in [Fig. 8\(d\)](#)] and stays in the evanescent field during cell migration (in [Visualization 1](#), it can be noticed for cells migrating between $+10 \text{ h}$ and $+17 \text{ h}$, between $+31 \text{ h}$ and $+37 \text{ h}$, and between $+47 \text{ h}$ and $+50 \text{ h}$). We attribute this dark part to the nucleus because this rather circular object has, as expected, a higher RI (~ 1.39) than the surrounding cytoplasm (~ 1.36). Note that when the nuclei are visible, they have the characteristic size of C2C12 myoblast nuclei, as observed with DIC ($20 \pm 4 \mu\text{m} \times 16 \pm 3 \mu\text{m}$), and blinking spots of submicrometer size appear ahead of the nuclei in the cell migration direction as the possible indication of focal adhesion sites [57].

If we compare SSPM images with former HRSPM images of fixed or living cells in a liquid medium ([43–49]), we can emphasize some drastic differences. Peterson *et al.*, [45] were the only ones to pinpoint focal adhesion sites (for fixed vascular smooth muscle cells A10 and human carcinoma cells HepG2) and extracellular materials (for fixed adenocarcinomic alveolar basal epithelial cells A549). Moh *et al.* [43] observed higher contrast domains corresponding to nuclei in fixed mouse fibroblast 3T3 cells. Jamil's *et al.* [46] were the first to reach the SSPM resolution on single filopodia of living cells. Since the cell substratum interaction depends on the cell type and the adhesion conditions (kind of surface, culture medium, and cell cycle phase), further tests on various cells and culture medium conditions should bring more insight on the cell substratum adhesion with HRSPM.

B. Adherence Evolution

During the 50 h recording, cells detaching or adhering to the surface can be dynamically observed in [Visualization 1](#). If we compare two images, one at the beginning [[Fig. 9\(a\)](#)] and the other one at the end [[Fig. 9\(b\)](#)] of the experiment, the image contrast is quite different, with a much lower proportion of bright regions in the latter than in the former. We have computed the normalized PDF of the two images [[Figs. 9\(c\)](#) and [9\(d\)](#)]. These histograms have a bimodal shape, with a first bump corresponding to the background noise and a second bump corresponding to the cells. These histograms can be fitted by the sum of two Gaussian distributions of parameters A_i, I_i, σ_i , with $i = 1, 2$, for each bump [green curves in [Figs. 9\(c\)](#) and [9\(d\)](#)]. A_i, I_i, σ_i are, respectively, the maximum amplitude, the mean value, and the standard deviation of each Gaussian curve. When comparing the rightmost bump corresponding to cell population, we notice a significant decrease of

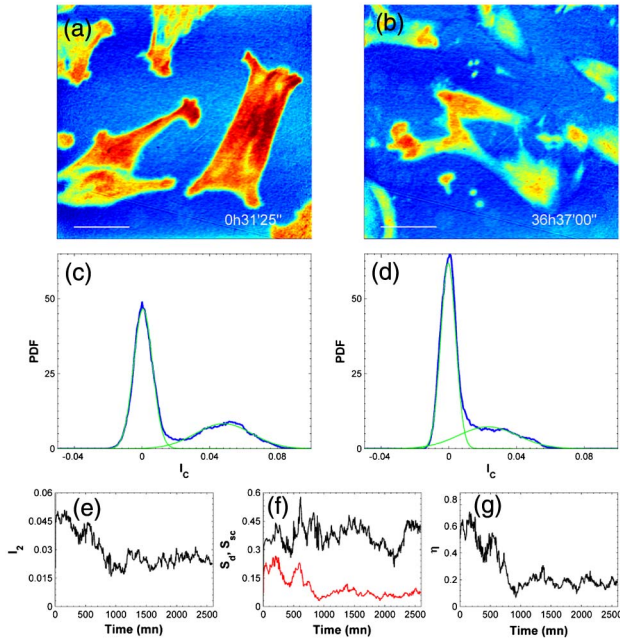


Fig. 9. Evolution in time of living cell adhesion. SSPM images: (a) at the beginning of Visualization 1 and (b) 36.5 h later at the end of recording. The scale bar is 20 μm and the intensity scale is [0.02; 0.08]. (c) and (d) are the corresponding PDF of images (a) and (b). Fits with a bimodal Gaussian distribution are plotted in green. (e) Corrected intensity $I_2(t)$ inside the cells versus time. (f) The global surface $S_d(t)$ covered by cells and probed by the SSPM (black) and the contact surface $S_{sc}(t)$ (red) versus time. (g) $\eta(t) = S_{sc}(t)/S_d(t)$ (see text).

I_2 from the beginning to the end of the time lapse recording, from 0.048 to 0.025, with a remarkable transition around 1000 min [Fig. 9(e)]. As probed by the SSPM, the global surface ratio covered by cells, $S_d = A_2\sigma_2/(A_1\sigma_1 + A_2\sigma_2)$ is more or less 40% and this value does not change much on average within the 50 h recording [Fig. 9(f)]. The decrease of I_2 without a significant change of S_d is an indication of some evolution in cell adhesion. Actually, the cells are in strong contact with the surface at the beginning of the recording and lose progressively this tight contact with time.

The surface proportion S_{sc} of the cells that are in tight contact with gold can be quantified. We delimit tight contact zones where the SSPM intensity is higher than half the maximum: $S_{sc} = \{(x, y)/I_c(x, y) > \max(I_c)/2\}/S_{\text{tot}}$, where S_{tot} is the total surface of the image. Two kinds of cell surface interaction fulfill this condition. First, cells that do not touch the surface but are close enough so that their bodies intersect the evanescent field [similar to the configuration in Fig. 7(c)]. This case is observed for rather spherical cells that are 60 nm or less away from the gold surface. Second, rather spreaded cells with rather thick parts (>60 nm) of their body that touch (tight contact) the gold surface. S_{sc} is plotted in red in Fig. 9(f): at the beginning of the recording, cells in strong contact cover about 25% of the image surface and this value decreases down to $\sim 5-10\%$ at the end of the experiment, with a transition at around 1000 min. Let us define $\eta = S_{sc}/S_d$, the ratio of the cell surface in strong contact S_{sc} and the cell surface detected by the SSPM.

As shown in Fig. 7(g), this ratio is important (60%) at the beginning but starts decreasing rather fast to reach $\sim 20\%$ after ~ 1000 min (3 fold decrease) and then stabilizes at the end of the recording. This modification of η points out a change in the cell adhesion, with a rather sharp transition at 1000 min. This change in adhesion comes along with some decrease of the number of observed filopodia and their almost total disappearance after ~ 1000 min (see Visualization 1).

This transition is certainly related to a modification of the experimental conditions inside the chamber. Since the chamber is sealed, we suspect that the pH value of the DMEM culture medium has decreased because of lack of CO_2 maintenance around 10%. Since cellular respiration produces carbon dioxide and lactate that can acidify the culture medium over time, this may significantly affect the microenvironment of the cells due to the limited volume of growth medium inside our sealed chambers (25 μl) [58]. This decrease of the culture medium pH might impact the intracellular pH and in turn limit the adhesion maturation by focal adhesion kinase (FAK) pathway [59]. Also, the amount of glucose may have decreased during the video recording, inducing a modification of the cell metabolic synthesis of ATP, resulting in a modification of the morphology, adhesion, and motility of the cells. Finally, during the cell cycle, the cells are synthesizing ECM proteins, cytokines, and cellular fragments that likely absorb on gold to form some hybrid layers with different adherence properties for the cells.

4. CONCLUSION

We report, to the best of our knowledge, the first high resolution surface plasmon imaging of living cells in a culture medium over long periods of time (50 h) that combines SSPM and a fibered interferometer. This SSPM does not suffer from aberrations and resolution loss faced by classical wide field microscopes. We have shown that the SSPM setup is suited for capturing the tightness of cell substrate contacts in real time and their detachment as well as their anchoring. It is also suited to characterize the local fluctuations of filopodia and lamellipodia protrusion shapes and velocities [60]. Besides its ability to capture the migration of adherent cells (few $\mu\text{m}/\text{minute}$), it can be used to compare cells with different strengths of adhesion in order to better understand their interactions with their ECM. Note that by replacing the piezoelectric stage by a motorized stage, a much wider area could be scanned, allowing us to follow long distance cell migration and interaction and to increase the size of the cell statistical sample on a single experiment. Even if scanning methods are limited for studying fast dynamics (faster than minutes), recent technical developments in beam focusing engineering [61] could be a future opportunity for improving the SSPM image resolution. Another possible improvement could be to couple SSPM with DIC and/or fluorescence microscopy in order to address deeper biological issues. In particular, SSPM could straightforwardly be extended to real time high resolution imaging of cell cell interactions in epithelial layers, since it was recently demonstrated by prism coupled SPR [62,63] that these layers likely play the role of wave guides. Indeed, as shown in a previous work [53], SSPM can perform submicron resolution imaging of wave guide

defects. This opens a new field of applications of this HRSPM method to thin tissue and multilayer characterization.

APPENDIX A

CELL CULTURE

C2C12 mouse myoblast cell line (American Type Culture Collection, Manassas, VA, number CRL 1772) was cultured in high glucose (4,5 g/L) Dulbecco's Modified Eagle Medium (DMEM, PAA, GE Healthcare Life Science, Munich, Germany) supplemented with 20% fetal bovine serum (FBS, PAA), 1% of penicillin streptomycin (10,000 U/mL, Gibco) and 10 mM HEPES. Myoblasts were kept in a 5% CO₂ atmosphere at 37°C on 90 mm petri dishes and fresh culture medium was replaced every 2 days. At 70% of confluence to avoid differentiation, dishes were washed twice with PBS and cells were dissociated from substrate using 0.25% trypsin EDTA (Gibco) for 3 min at 37°C. Then cells were replated at a lower concentration (30% of confluence). Fresh culture medium was replaced each 2–3 days. Cells were used at the 12th passage.

SAMPLE PREPARATION

High RI (1.648) coverslips were coated with a 1.5 nm chromium layer and a 45 nm gold layer by cathodic sputtering. 2.5×10^4 cells were seeded on the gold surface within their culture medium and kept 4 h at 37°C with 5% CO₂ to let them adhere on the gold substrate. The gold coated coverslip was then sealed to a second glass coverslip with a double face sticky spacer delimiting a rectangular chamber filled with the culture medium (Gene Frame from Thermo Fisher Scientific). To enable long term image acquisition with motionless biological objects, adherent myoblasts were fixed with 4% paraformaldehyde (PFA, Sigma) for 20 min at room temperature after rinsing the sample twice with PBS. Fixed myoblasts were covered with 25 µl of phosphate buffered saline (PBS) while living myoblasts were maintained in 25 µl of culture medium. The dimension and the volume of the chamber are 1 cm × 1 cm × 250 µm and 25 µl. The time interval of 65 s between two images includes 10 s for $V(z)$ recording, 50 s for image acquisition, and a 5 s additional delay.

$|V(z)|$ COMPUTATIONS

The theoretical $|V(z)|$ curves were computed from the integral of the reflectivities $R_{p,s}(\theta)$ [Eq. (1)]. These reflectivities R_p and R_s were obtained from Fresnel equations for a 4 layer model comprising (i) a semi infinite coupling medium (1.57 NA Zeiss objective lens, High RI coverslip, and immersion oil) with RI $n_1 = 1.648$, (ii) a 45 nm gold layer with a complex dielectric coefficient $\epsilon_2(632.8 \text{ nm}) = 12.31 + 1.22i$, (iii) a dielectric layer with thickness h and refractive index n_3 , and (iv) a semi infinite liquid medium with refractive index $n_4 = 1.33$.

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